The Comprehensive Table of Contents Headings and Hierarchy

Revision History

Date	Version	Summary of Changes	
2004-07	1.0	Original version	
2005-06-16	1.1	Corrections and additions to the mapping tables	
2005-07-06 1.2 Corrections to the headings		Corrections to the headings	
2012-06-01	2.0	Corrections and additions to the mapping tables based on major	
		update to Module 1 specifications (<u>Summary of Changes in Section</u>	
		C of Appendix 2)	
2012-11-01	2.1	Modified the heading for 1.16 and added REMS and non-REMS	
		sub-headings (Summary of Changes in Section B of Appendix 2)	
2013-08-23	2.2	Added two new attributes for 1.15.2.1 (Summary of Changes in	
		Section A of Appendix 2)	
2014-02-07	2.3	Modified the heading for 1.15.1.5 (<u>Summary of Changes in Section</u>	
		A of Appendix 2)	

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- 3.2.S.2.5 Process Validation and/or Evaluation
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- 3.2.S.7.2 Post Approval Stability Protocol and Stability Commitment
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3.2.P.1 Description and composition of the drug product

3.2.P.2 Pharmaceutical development

3.2.P.3 Manufacture

- 3.2.P.3.1 Manufacturer(s)
- 3.2.P.3.2 Batch Formula
- 3.2.P.3.3 Description of Manufacturing Process and Process Controls
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Documentation of inter laboratory standardization methods of

quality assurance procedures if used

Publications based on the study

Important publications referenced in the report

Compliance and/or drug concentration data

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Data tabulation datasets

Data definitions

Data listing datasets

Data listing datasets

Data definitions

Analysis datasets

Analysis datasets

Analysis programs

Data definitions

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See Primary pharmacodynamics Study report and related information for headings

4.2.1.3 Safety pharmacology

Study report [identification number] and related information

See Primary pharmacodynamics Study report and related information for headings

4.2.1.4 Pharmacodynamic drug interactions

Study report [identification number] and related information

See Primary pharmacodynamics Study report and related information for headings

4.2.2 Pharmacokinetics

4.2.2.1 Analytical methods and validation reports

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See Primary pharmacodynamics Study report and related information for headings

4.2.2.2 Absorption

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See Primary pharmacodynamics Study report and related information for headings

4.2.2.3 Distribution

Study report [identification number] and related information

See Primary pharmacodynamics Study report and related information for headings

4.2.2.4 Metabolism

Study report [identification number] and related information

See Primary pharmacodynamics Study report and related information for headings

4.2.2.5 Excretion

Study report [identification number] and related information

See Primary pharmacodynamics Study report and related information for headings

4.2.2.6 Pharmacokinetic drug interactions

Study report [identification number] and related information

See Primary pharmacodynamics Study report and related information for heading Statement of QA differences

4.2.2.7 Other pharmacokinetic studies

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See Primary pharmacodynamics Study report and related information for headings

4.2.3 Toxicology

4.2.3.1 Single dose toxicity [Species and route]

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See Primary pharmacodynamics Study report and related information for headings

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Study report [identification number] and related information

See Primary pharmacodynamics Study report and related information for headings

4.2.3.3 Genotoxicity

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See Primary pharmacodynamics Study report and related information for headings

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See Primary pharmacodynamics Study report and related information for headings

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See Primary pharmacodynamics Study report and related information for headings

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See Primary pharmacodynamics Study report and related information for headings

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Data listing datasets

Data definitions

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Analysis datasets

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Data definitions

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Study report [identification] and related information

See example under bioavailability (BA) Study reports and related information for headings

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Study report [identification] and related information

See example under bioavailability (BA) Study reports and related information for headings

5.3.2.3 Reports of studies using other human biomaterials

Study report [identification] and related information

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5.3.3 Reports of human pharmacokinetic (PK) studies

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Study report [identification] and related information

See example under bioavailability (BA) Study reports and related information for headings

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See example under bioavailability (BA) Study reports and related information for headings

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Study report [identification] and related information

See example under bioavailability (BA) Study reports and related information for headings

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See example under bioavailability (BA) Study reports and related information for headings

5.3.4.2 Patient PD and PK/PD Study reports and related information

Study report [identification] and related information

See example under bioavailability (BA) Study reports and related information for headings

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5.3.5.2 Study reports and related information of uncontrolled clinical studies

Study report [identification] and related information

See example under bioavailability (BA) Study reports and related information for headings

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Integrated summary of efficacy report

Analysis datasets

Analysis programs

5.3.5.4 Other Study reports and related information

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Antiviral reports

5.3.6 Reports of postmarketing experience

Postmarketing periodic adverse event drug experience report description

5.4 Literature references

Appendix I – Mapping Section *IND*

CFR	Citation/Source		CTD /*STF Heading/**Attribute(s)		
NUMBER	TITLE	MODULE	NUMBER	TITLE	
312.23(a)(1)	Cover sheet (Form FDA–1571)	1	1.1	**Forms form-type=1571	
FDAAA	Certification of compliance:	1	1.1	**Forms form-type=3674	
	Form FDA 3674				
BsUFA	Form FDA 3792: Biosimilar User	1	1.1	**Forms form-type=3792	
	Fee Cover Sheet				
312.31(b)(1)	Statement of the nature and	1	1.2	Cover letters	
	purpose of the information				
	amendment				
	Change of address or corporate	1	1.3.1.1	Change of address or corporate name	
	name				
	NOTE: Includes DMF original				
	address or corporate name or				
	change in DMF address or				
	corporate name				
	Change in contact/agent	1	1.3.1.2	Change in contact/agent	
	NOTE: Includes DMF original				
	contact/agent or change in DMF				
	contact/agent				
	Change in ownership	1	1.3.1.3	Change in sponsor	
312.52	Transfer of obligations to a	1	1.3.1.4	Transfer of obligation	
	contract research organization				
312.22(d)	General principles of the IND		1.4.1	Letter of authorization	
	submission				
312.23(b)	Written statement of	1	1.4.2	Statement of right of reference	
	authorization for references				
	(copy of LOA received from				
	DMF holders - submitted by				
	BLA, NDA, or IND applicants)				

CFR Citation/Source			CTD /*STF Heading/**Attribute(s)			
NUMBER	TITLE	MODULE	NUMBER	TITLE		
312.23(b)	Information previously	1	1.4.4	Cross-reference to previously submitted		
312.23(a)(3)(ii)	submitted			information		
312.38	Withdrawal of an IND	1	1.5.1	Withdrawal of an IND		
312.45(a)	Request for Inactive status	1	1.5.2	Inactivation request		
312.45(d)	Request to resume clinical investigation under an inactive IND	1	1.5.3	Reactivation request		
	Reinstatement request	1	1.5.4	Reinstatement request		
312.47	Meeting request	1	1.6.1	Meeting request		
PDUFA Agreements						
312.47	Meeting background material	1	1.6.2	Meeting background materials		
PDUFA Agreements						
312.47	Correspondence regarding a	1	1.6.3	Correspondence regarding meetings		
PDUFA Agreements	meeting					
FDAMA	Fast track designation request	1	1.7.1	Fast track designation request		
FDAMA	Fast track designation	1	1.7.2	Fast track designation withdrawal request		
	withdrawal request					
FDAMA	Rolling review request	1	1.7.3	Rolling review request		
FDAMA	Correspondence regarding fast	1	1.7.4	Correspondence regarding fast track/rolling		
	track/rolling review			review		
FDAMA	Special protocol assessment	1	1.8.1	Clinical study		
	request: clinical study					
PDUFA Agreements	Special protocol assessment	1	1.8.2	Carcinogenicity study		
	request: carcinogenicity study					
PDUFA Agreements	Special protocol assessment	1	1.8.3	Stability study		
	request: stability study					
	Animal efficacy study for	1	1.8.4.	Animal efficacy study for approval under		
	approval under the animal rule			the animal rule		
PREA	Request for waiver of pediatric	1	1.9.1	Request for waiver of pediatric studies		
312.47(b)(1)(iv)	studies					

CFR (Citation/Source		CTD /*ST	F Heading/**Attribute(s)
NUMBER	TITLE	MODULE	NUMBER	TITLE
PREA	Request for deferral of pediatric	1	1.9.2	Request for deferral of pediatric studies
312.82	studies			
312.47(b)(1)(iv)				
BPCA	Proposed pediatric study request and amendments	1	1.9.4	Proposed pediatric study request and amendments
PREA	Correspondence regarding	1	1.9.6	Other correspondence regarding pediatric
BPCA	pediatric exclusivity or PREA requirements			exclusivity or study plans
312.48	Scientific and medical disputes	1	1.10.1	Request for dispute resolution
312.48	Scientific and medical disputes	1	1.10.2	Correspondence related to dispute resolution
312.31	Information amendment:	1	1.11.1	Quality information amendment
	Chemistry - information not covered under Module 3			
312.31	Information amendment:	1	1.11.2	Nonclinical information amendment
	Toxicology - information not covered under Module 4			
312.31	Information amendment: Clinical - information not covered under Module 5	1	1.11.3	Clinical information amendment
312.31	Multiple Information amendment	1	1.11.4	Multiple module information amendment
312.82(a)	Pre-IND correspondence	1	1.12.1	Pre-IND correspondence
312.8(b)	Charging for investigational drugs under an IND	1	1.12.2	Request to charge for clinical trial
312.8(c)	Charging for investigational drugs under an IND	1	1.12.3	Request to charge for expanded access
312.31(b)(3)	Request for comment on information amendment	1	1.12.4	Request for comments and advice
312.41	Comment and advice on an IND	1	1.12.4	Request for comments and advice
312.10	Waivers (including PSUR waiver)	1	1.12.5	Request for a waiver

CFR (Citation/Source		CTD /*ST	F Heading/**Attribute(s)
NUMBER	TITLE	MODULE	NUMBER	TITLE
312.54	Exception from informed	1	1.12.6	Exception from informed consent for
	consent for research			emergency research
312.54	Public disclosure – exception	1	1.12.7	Public disclosure statement for exception
	from informed consent for			from informed consent for emergency
	research			research
312.54	IRB disapproval of exception	1	1.12.8	Correspondence regarding exception from
	from informed consent for			informed consent for emergency research
	research			
312.31(a)(2)	Report regarding the	1	1.12.9	Notification of discontinuation of clinical
	discontinuation of a clinical			trial
212.22(a)(7)(iv)(a)	investigation	1	1.12.14	Environmental analysis
312.23(a)(7)(iv)(e)	Environmental analysis requirements	1	1.12.14	Environmental analysis
316 Subpart C	Orphan Drug	1	1.12.17	Orphan drug designation
312.33(b)(6)	Annual Report: A list of	1	1.13.1	Summary of nonclinical studies
312.33(0)(0)	preclinical studies	1	1.13.1	Summary of nonclinical studies
312.33(b)(5)	Annual Report: A brief	1	1.13.2	Summary of clinical pharmacology
312.00(0)(0)	description of the drug's	_	1.12.2	information
	actions			
312.33(b)(1)	Annual Report: A narrative or	1	1.13.3	Summary of safety information
	tabular summary showing the			
	most frequent and most serious			
	adverse experiences by the body			
	system			
312.33(b)(2)	Annual Report: A summary of	1	1.13.3	Summary of safety information
	all IND safety reports			
312.33(b)(3)	Annual Report: A list of	1	1.13.3	Summary of safety information
	subjects who died			
312.33(b)(4)	Annual Report: A list of	1	1.13.3	Summary of safety information
	subjects who dropped out			

CFR (Citation/Source		CTD /*STF Heading/**Attribute(s)		
NUMBER	TITLE	MODULE	NUMBER	TITLE	
312.33(b)(7)	Annual Report: A summary of any significant manufacturing changes	1	1.13.5	Summary of manufacturing changes	
312.33(b)(7)	Annual Report: A summary of any significant microbiological changes	1	1.13.6	Summary of microbiological changes	
312.33(a)	Annual report individual study information	1	1.13.8	Individual study information	
312.33(c)	Annual Report: A description of the general investigational plan	1	1.13.9	General investigational plan	
312.33(f)	Annual Report: A brief summary of significant foreign marketing developments	1	1.13.10	Foreign marketing	
312.33(g)	Annual Report: Log of outstanding business(optional)	1	1.13.14	Log of outstanding regulatory business	
	Development safety update report (DSUR)	1	1.13.15	Development safety update report (DSUR)	
312.6	Draft labeling text	1	1.14.1.3	Draft labeling text	
	Label comprehension studies	1	1.14.1.4	Label comprehension studies	
312.23(a)(5)	Investigator brochure	1	1.14.4.1	Investigator brochure	
312.33(d)	Annual Report: Investigators brochure	1	1.14.4.1	Investigator brochure	
312.23(a)(7)(iv)(d)	Labeling	1	1.14.4.2	Investigational drug labeling	
	Foreign labeling	1	1.14.5	Foreign labeling	
	Proprietary names	1	1.18	Proprietary names	
Project BioShield Act of 2004	Emergency Use Authorization	1	1.19	Pre-EUA and EUA	
312.23(a)(3)(iv)	A brief description of the overall plan	1	1.20	General investigational plan for initial IND	

CFR	Citation/Source		CTD /*ST	F Heading/**Attribute(s)
NUMBER	TITLE	MODULE	NUMBER	TITLE
312.23(a)(3)(i)	Introductory statement	2	2.2	Introduction to summary
312.23(a)(7)(a), (b)	Chemistry, manufacturing, and	2	2.3	Quality overall summary
and (c)	controls			
312.23(a)(8)	Pharmacology and toxicology information	2	2.4	Nonclinical overview
312.23(a)(9)	Previous human experience	2	2.5	Clinical overview
312.23(a)(3)(ii-iii)	Introductory statement	2	2.5	Clinical overall summary
312.23(a)(8)	Pharmacology and toxicology	2	2.6	Nonclinical written and tabulated summaries
	information			[use appropriate sections]
312.23(a)(9)	Previous human experience	2	2.7	Clinical summary [use appropriate sections]
312.23(a)(10)(i)	Drug dependence and abuse	2	2.7.4	Summary of Clinical Safety
312.23(a)(8)	Pharmacology and toxicology information	4	4.2	Study reports [use appropriate sections]
312.23(a)(9)	Previous human experience	5	5.3	Clinical study reports and related information [use appropriate sections]
312.30(a)	New protocol	5	5.3	Protocol [under specific study]
312.30(b)	Changes in protocol	5	5.3	Protocol [under specific study]
312.30(c)	New investigator	5	5.3	List and description of investigators and sites [under specific study]
312.23(a)(6)	Protocol	5	5.3	*Protocol [under specific study]
312.32	IND safety reports	5	5.3	*IND safety report [under specific study]
312.33(e)	Annual Report: A description of	5	5.3	*Protocol [under the specific study]
	any significant Phase 1 protocol			
	modifications made during the			
212 220	previous years and	-		100
312.320	Treatment protocol	5	5.3	*Protocol [under specific study]
312.120(b)(1)	Foreign clinical studies not	5	5.3	*List and description of investigators and
	conducted under the IND:			sites [under specific study]
	Investigator's qualification			

CFR (Citation/Source		CTD /*ST	TF Heading/**Attribute(s)
NUMBER	TITLE	MODULE	NUMBER	TITLE
312.120(b)(2)	Foreign clinical studies not conducted under the IND: Research facility	5	5.3	*List and description of investigators and sites [under specific study]
312.120(b)(3)	Foreign clinical studies not conducted under the IND: Detailed summary	5	5.3	Use appropriate sections [under specific study]
312.120(a)(1)	Foreign clinical studies not conducted under the IND: Conformance with ethical principles	5	5.3	*List of IECs or IRBs and consent forms [under specific study]
312.23(a)(11)	Relevant information	1, 2, 3, 4, or 5	As needed	Use appropriate sections
312.23(c)	Material in a foreign language (English translations)	1, 2, 3, 4, or 5	As needed	Use appropriate sections
312.23(a)(10)(iv)	Other information	2, 3, 4, or 5	As needed	Use appropriate sections
312.23(a)(10)(ii)	Radioactive drugs	2, 4, or 5	As needed	Use appropriate sections
312.23(a)(7)(a), (b) and (c)	Chemistry, manufacturing and controls	3	As needed	Quality [use appropriate sections]
312.31(a)(1),	Information amendment: Chemistry	3	As needed	Use appropriate sections
312.120(b)(4)	Foreign clinical studies not conducted under the IND: A description of the drug substance and drug product	3	As needed	Use appropriate sections
312.31	Information amendment: Toxicology	4	As needed	Use appropriate sections
312.31	Information amendment: Clinical	5	As needed	Use appropriate sections
312.23(a)(2)	Table of contents	N/A	N/A	N/A

NDA and BLA

CFR	Citation/Source		CTD /*ST	TF Heading/**Attribute(s)
NUMBER	TITLE	MODULE	NUMBER	TITLE
314.50(a)	Application Form FDA 356h	1	1.1	**Forms form-type=356h
601.2				
PDUFA	User fee cover sheet: Form FDA 3397	1	1.1	**Forms form-type=3397
BsUFA	Form FDA 3792: Biosimilar User Fee Cover Sheet	1	1.1	**Forms form-type=3392
314.81(b)(2)	Annual report transmittal: Form FDA 2252	1	1.1	**Forms form-type=2252
314.81(b)(3)(i) 601.12(f)(4)	Transmittal of advertisements and promotional labeling: Form FDA 2253	1	1.1	**Forms form-type=2253
601.12 (f)	Transmittal of labels and circulars: Form FDA 2567	1	1.1	**Forms form-type=2567
	Cover letters	1	1.2	Cover letters
	Change of address or corporate name NOTE: Includes DMF original address or corporate name or change in DMF address or corporate name	1	1.3.1.1	Change of address or corporate name
	Change in contact/agent NOTE: Includes DMF original contact/agent or change in DMF contact/agent	1	1.3.1.2	Change in contact/agent
314.50(d)(5)(x)	Transfer of obligations to CRO	1	1.3.1.4	Transfer of obligation
314.72 601.4	Change in ownership of an application	1	1.3.1.5	Change in ownership of an application or reissuance of license
314.50(d)(1)(v)	Field copy certification	1	1.3.2	Field copy certification
GDEA	Debarment certification	1	1.3.3	Debarment certification

CFR Citation/Source			CTD /*S7	FF Heading/**Attribute(s)
NUMBER	TITLE	MODULE	NUMBER	TITLE
314.50(k)	Financial certification and	1	1.3.4	Financial certification and disclosure
601.2(a)	disclosure statement (Form FDA			
	3454 and Form FDA 3455)			
314.50(h)	Patent Information (Form FDA	1	1.3.5.1	Patent information
314.53(e)	3542a and Form FDA 3542)			
314.50(i)	Patent certification	1	1.3.5.2	Patent certification
314.52(e)				
314.50(j)	Claimed exclusivity	1	1.3.5.3	Exclusivity claim
FDAAA	Tropical disease priority review	1	1.3.6	Tropical disease priority review voucher
	voucher			
314.420(d)	Incorporating DMF information	1	1.4.1	Letter of authorization
	by reference (authorization from			
	DMF holder)			
314.50(g)(1)	Written statement of	1	1.4.2	Statement of right of reference
	authorization for references			
	(copy of LOA received from			
	DMF holders - submitted by			
	BLA, NDA, or IND applicants)			
314.420(d)	List of authorized persons to	1	1.4.3	List of authorized persons to incorporate by
	incorporate by reference			reference
314.50(g)(1)	Reference to information	1	1.4.4	Cross-reference to previously submitted
	previously submitted			information
314.65	Withdrawal of an unapproved	1	1.5.5	Withdrawal of an unapproved NDA, ANDA
	application			or Supplement
314.50	Withdrawal of listed drug	1	1.5.6	Withdrawal of listed drug
314.150(c)	Withdrawal of approval	1	1.5.7	Withdrawal of approval of an application or
				revocation of license
314.150	Withdrawal of approval by the	1	1.5.7	Withdrawal of approval of an application or
601.5	FDA			revocation of license
314.102	Communications:	1	1.6.1	Meeting request
	Meetings			

CFR	Citation/Source		CTD /*S7	TF Heading/**Attribute(s)
NUMBER	TITLE	MODULE	NUMBER	TITLE
314.102	Communications:	1	1.6.2	Meeting background materials
	Meetings			
314.102	Communications:	1	1.6.3	Correspondence regarding meetings
	Meetings			
FDAMA	Fast track designation request	1	1.7.1	Fast track designation request
FDAMA	Fast track designation	1	1.7.2	Fast track designation withdrawal request
	withdrawal request			
FDAMA	Rolling review request	1	1.7.3	Rolling review request
FDAMA	Correspondence regarding fast	1	1.7.4	Correspondence regarding fast track/rolling
	track/rolling review			review
PREA	Request for waiver of pediatric	1	1.9.1	Request for waiver of pediatric studies
314.55(c)	studies			
601.27(c)				
PREA	Request for deferral of pediatric	1	1.9.2	Request for deferral of pediatric studies
314.55(b)	studies			
601.27(b)				
BPCA	Request for pediatric exclusivity	1	1.9.3	Request for pediatric exclusivity
	determination/Form FDA 3437			determination
BPCA	Proposed pediatric study request	1	1.9.4	Proposed pediatric study request and
	and amendments			amendments
PREA	Correspondence regarding	1	1.9.6	Other correspondence regarding pediatric
BPCA	pediatric exclusivity or PREA			exclusivity or study plans
	requirements			
314.103(c)	Scientific and medical disputes	1	1.10.1	Request for dispute resolution
314.103(c)	Scientific and medical disputes	1	1.10.2	Correspondence related to dispute resolution
314.60	Amendment to an unapproved	1	1.11.1	Quality information amendment
	application: Chemistry			
	(information not covered under			
	Module 3)			
314.60	Amendment to an unapproved	1	1.11.2	Nonclinical information amendment
	application: Toxicology			

CFR Citation/Source		CTD /*STF Heading/**Attribute(s)		
NUMBER	TITLE	MODULE	NUMBER	TITLE
	(information not covered under Module 4)			
314.60	Amendment to an unapproved application: Clinical (information not covered under Module 5)	1	1.11.3	Clinical information amendment
314.60	Multiple information amendment:	1	1.11.4	Multiple module information amendment
	Request for comment and advice	1	1.12.4	Request for comments and advice
314.90 600.90	Waivers (including PSUR waiver)	1	1.12.5	Request for a waiver
GDEA	Generic drug enforcement act statement	1	1.12.10	Generic drug enforcement act statement
314.50(d)(1)(iii) 601.2	Environmental impact	1	1.12.14	Environmental analysis
320.22 (a)	Request for waiver of in vivo bioavailability studies	1	1.12.15	Request for waiver of in vivo bioavailability studies
314.81(b)(1)	Field alert reports	1	1.12.16	Field alert reports
316 Subpart C	Orphan drug	1	1.12.17	Orphan drug designation
314.81(b)(2)(i) 601.12(d)	Annual Report: Summary	1	1.13.1	Summary of nonclinical studies
314.81(b)(2)(i) 601.12(d)	Annual Report: Summary	1	1.13.2	Summary of clinical pharmacology information
314.81(b)(2)(i) 601.12(d)	Annual Report: Summary	1	1.13.3	Summary of safety information
314.81(b)(2)(i) 601.12(f)(3)	Annual Report: Summary	1	1.13.4	Summary of labeling changes
314.81(b)(2)(i) 601.12(d)	Annual Report: Summary	1	1.13.5	Summary of manufacturing changes
314.81(b)(2)(i) 601.12(d)	Annual Report: Summary	1	1.13.6	Summary of microbiological changes
314.81(b)(2)(i)	Annual Report: Summary	1	1.13.7	Summary of other significant new

CFR Citation/Source		CTD /*STF Heading/**Attribute(s)		
NUMBER	TITLE	MODULE	NUMBER	TITLE
601.12(d)				information
314.81(b)(2)(ii)	Annual Report: Distribution data	1	1.13.11	Distribution data
314.81(b)(2)(vii)	Annual Report: Status report of	1	1.13.12	Status of postmarketing study commitments
601.70	clinical and nonclinical			and requirements
	toxicology postmarketing study			
	commitments			
314.81(b)(2)(viii)	Status report of other (chemistry,	1	1.13.13	Status of other postmarketing studies and
	manufacturing, controls)			requirements
	postmarketing study			
214.01(1)(2)(;)	commitments	1	1 10 14	T C
314.81(b)(2)(ix)	Annual Report: Log of	1	1.13.14	Log of outstanding regulatory business
214.50(-)(2)('')	outstanding regulatory business	1	1 1 4	
314.50(e)(2)(ii) 601.14	Copies of the labeling and all	1	1.14	Use appropriate sections
	labeling for the drug product Annual Report: Labeling	1	1.14	Has appropriate sections
314.81(b)(2)(iii) 601.14(f)(3)	Annual Report: Labering	1	1.14	Use appropriate sections
314.50	Draft carton and container labels	1	1.14.1.1	Draft carton and container labels
601.14	Dian canon and container labers	1	1.14.1.1	Diant carton and container labers
314.50(c)(2)(i)	The proposed text of the labeling	1	1.14.1.2	Annotated draft labeling text
311.30(0)(2)(1)	with annotations	1	1.1 1.1.2	Timotated draft labeling text
314.50(e)(2)(ii)	Draft labeling text	1	1.14.1.3	Draft labeling text
601.2 601.14		_		
	Label comprehension studies	1	1.14.1.4	Label comprehension studies
	Labeling history	1	1.14.1.5	Labeling history
314.50(e)(2)(ii)	Final carton or container labels	1	1.14.2.1	Final carton or container labels
601.2				
314.50(e)(2)(ii)	Final package insert (package	1	1.14.2.2	Final package insert (package inserts, patient
601.2; 601.14	inserts, patient information,			information, medication guides)
	medication guides)			
314.50(e)(2)(ii)	Final labeling text	1	1.14.2.3	Final labeling text
601.2; 601.14				

CFR Citation/Source			CTD /*STF Heading/**Attribute(s)		
NUMBER	TITLE	MODULE	NUMBER	TITLE	
	Foreign labeling	1	1.14.5	Foreign labeling	
314.81(b)(3)(i)	Product labeling for 2253	1	1.14.6	Product labeling for 2253 submissions	
601.12(f)(4)	submissions (if applicable)				
314.81(b)(3)(i)	Regulations related to	1	1.15	Promotional material **[promotional-	
601.12(f)(4)	promotional materials [use			material-audience-type]	
314.550	appropriate sections]				
601.45					
202.1(j)(4)					
314.640					
601.94					
202.1					
202.1(j)(4)	Request for advisory comments	1	1.15.1.1	Request for advisory comments on launch	
	on launch materials			materials	
202.1(j)(4)	Request for advisory comments	1	1.15.1.2	Request for advisory comments on non-	
	on non-launch materials			launch materials	
314.550	Presubmission of launch	1	1.15.1.3	Presubmission of launch promotional	
601.45	promotional materials for			materials for accelerated approval products	
	accelerated approval of products				
	for serious or life-threatening				
	illnesses				
314.640	Presubmission of launch	1	1.15.1.3	Presubmission of launch promotional	
601.94	promotional materials for			materials for accelerated approval products	
	products approved when human				
	efficacy studies are not ethical or				
	feasible				
314.550	Presubmission of non-launch	1	1.15.1.4	Presubmission of non-launch promotional	
601.45	promotional materials for			materials for accelerated approval products	
	accelerated approval of products				
	for serious or life-threatening				
	illnesses				
314.640	Presubmission of non-launch	1	1.15.1.4	Presubmission of non-launch promotional	

CFR Citation/Source		CTD /*STF Heading/**Attribute(s)		
NUMBER	TITLE	MODULE	NUMBER	TITLE
601.94	promotional materials for products approved when human efficacy studies are not ethical or feasible			materials for accelerated approval products
202.1 Section 503C of the Food, Drug, and Cosmetic Act	Pre-dissemination review of television ads	1	1.15.1.5	Pre-dissemination review of television ads
202.1	Response to untitled letter or warning letter	1	1.15.1.6	Response to untitled letter or warning letter
202.1	Response to information request	1	1.15.1.7	Response to information request
202.1 314.81(b)(3)(i) 601.12(f)(4) 202.1(j)(4) 314.550 601.45 314.640 601.94	Correspondence accompanying materials previously missing or rejected	1	1.15.1.8	Correspondence accompanying materials previously missing or rejected
202.1 314.81(b)(3)(i) 601.12(f)(4) 202.1(j)(4) 314.550 601.45 314.640 601.94	Withdrawal request	1	1.15.1.9	Withdrawal request
202.1 202.1(j)(4) 314.550 601.45	Submission of annotated references	1	1.15.1.10	Submission of annotated references

CFR Citation/Source		CTD /*STF Heading/**Attribute(s)		
NUMBER	TITLE	MODULE	NUMBER	TITLE
314.640				
601.94				
202.1	General correspondence	1	1.15.1.11	General correspondence
314.81(b)(3)(i)	Regulations related to	1	1.15.2	Materials ** [promotional-material-doc-
601.12(f)(4)	promotional materials [use			type]
202.1(j)(4)	appropriate sections]			
314.550				
601.45				
314.640				
601.94				
202.1				
314.81(b)(3)(i)	Regulations related to	1	1.15.2.1	Material **[promotional-material-type,
601.12(f)(4)	promotional materials [use			material-id, issue-date]
202.1(j)(4)	appropriate sections]			
314.550				
601.45				
314.640				
601.94				
202.1				
202.1	Clean version	1	1.15.2.1.1	Clean version
314.81(b)(3)(i)				
601.12(f)(4)				
202.1(j)(4)				
314.550				
601.45				
314.640				
601.94				
202.1(j)(4)	Annotated version	1	1.15.2.1.2	Annotated version
314.550				
601.45				
314.640				

CFR	CTD /*STF Heading/**Attribute(s)			
NUMBER	TITLE	MODULE	NUMBER	TITLE
601.94				
202.1				
202.1(j)(4)	Annotated labeling version	1	1.15.2.1.3	Annotated labeling version
314.550				
601.45				
314.640				
601.94				
202.1				
202.1(j)(4)	Annotated references	1	1.15.2.1.4	Annotated references
314.550				
601.45				
314.640				
601.94				
202.1				
FDAAA 505-1	Risk evaluation and mitigation	1	1.16	Use the appropriate sections
[355-1]	strategies (REMS)			
FDAAA	Correspondence regarding	1	1.17.1	Correspondence regarding postmarketing
	postmarketing commitments			commitments
FDAAA	Correspondence regarding	1	1.17.2	Correspondence regarding postmarketing
	postmarketing requirements			requirements
	Proprietary names	1	1.18	Proprietary names
314.50(d)(5)(viii)	An integrated summary of the	2	2.5	Use appropriate sections
	benefits and risks			
314.50(c)(2)(ii) to	Summaries	2	As needed	Use the appropriate sections
(ix)				
314.50(d)(7)	Pediatric use section	2 and 5	As needed	Use appropriate sections
314.50(d)(1)(i) and	Chemistry, manufacturing and	3	As needed	Use the appropriate sections
(ii)	controls			
314.50(e)(2)(i)	Analytical methods	3	As needed	Use appropriate sections

CFR Citation/Source			CTD /*STF Heading/**Attribute(s)		
NUMBER	TITLE	MODULE	NUMBER	TITLE	
314.60	Amendment to an unapproved application: Chemistry	3	As needed	Use appropriate sections	
600.81	Distribution reports	3	3.2.R	Regional Information	
314.81(b)(2)(iv)	Annual Report: Chemistry, manufacturing, and controls	3	As needed	Use appropriate sections	
314.50(d)(2)	Nonclinical pharmacological and toxicology section	4	As needed	Use appropriate sections	
314.81(b)(2)(v)	Annual Report: Nonclinical laboratory studies	4	As needed	Use appropriate sections	
314.60	Amendment to an unapproved application: Toxicology	4	As needed	Use appropriate sections	
314.50(d)(5)(ix)	Statement of compliance with informed consent	5	5.3	*List of IECs or IRBs and consent forms [under specific study]	
314.50(d)(5)(xi)	Audited studies	5	5.3	*Audit certificates and reports [under specific study]	
314.50(d)(6)(i) and (ii)	Description of statistical analysis	5	5.3	*Documentation of statistical methods and interim analysis plans [under specific study]	
314.50(f)(1)	Case report tabulations	5	5.3	*Case report tabulations [use the appropriate sections under the specific study]	
314.50(f)(2)	Case report forms	5	5.3	*Case report forms [under the appropriate site and specific study]	
314.50(d)(5)(i) to (iv)	Clinical data section	5	5.3	Use appropriate sections	
314.50(d)(3)	Human pharmacokinetics and bioavailability sections	5	5.3	Use appropriate sections	
314.50(d)(5)(vii)	Potential for abuse	5	5.3	Use appropriate sections	
314.50(d)(5)(v)	An integrated summary of efficacy	5	5.3.4	Reports of analysis of data from more than one study [Use appropriate sections in integrated summary of efficacy STF]	
314.50(d)(5)(vi)(a)	An integrated summary of safety	5	5.3.4	Reports of analysis of data from more than one study [Use appropriate sections in integrated summary of safety STF]	

NDA and BLA Mapping Section

CFR Citation/Source		CTD /*STF Heading/**Attribute(s)		
NUMBER	TITLE	MODULE	NUMBER	TITLE
314.50(d)(5)(vi)(b)	Safety Update	5	5.3.5	Reports of analysis of data from more than
				one study [Use appropriate sections in
				integrated summary of safety STF]
314.50(d)(4)	Microbiology	5	5.3.5.4	Other study reports and related information
				[Use appropriate sections in microbiology
				STF]
314.80(c)(2)(ii)(a)	Periodic adverse drug experience	5	5.3.6	Postmarketing periodic adverse event drug
314.80(c)(2)(ii)(c)	 narrative summary and history 			experience report description
600.80(c)(20(ii)(A)	of actions			
600.80(c)(2)(ii)(C)				
314.70 and 314.71	Supplements and other changes	1, 2, 3, 4, 5	As needed	Use the appropriate sections
601.12	to approved applications			
314.420(a)	Drug master files	1, 2, 3, 4, 5	As needed	Use appropriate sections
314.60	Amendment to an unapproved	5	As needed	Use appropriate sections
	application: Clinical			
314.81(b)(2)(vi)	Annual Report: Clinical data	5	As needed	Use appropriate sections
315.50(b)	Index	N/A	N/A	N/A

ANDA

CFF	R Citation/Source		CTD /*STF	Heading/**Attribute(s)
NUMBER	TITLE	MODULE	NUMBER	TITLE
314.94(a)(1)	Application Form FDA 356h	1	1.1	**Forms form-type=356h
GDUFA	Form FDA 3794: Generic Drug User Fee Cover Sheet	1	1.1	**Forms form-type=3794
FDAAA	Certification of compliance: Form FDA 3674	1	1.1	**Forms form-type=3674
	Transmittal of labels and circulars: Form FDA 2567	1	1.1	**Forms form-type=2567
314.81(b)(3)(i)	Transmittal of advertisements and promotional labeling: Form FDA 2253	1	1.1	**Forms form-type=2253
	Cover letters	1	1.2	Cover letters
	Change of address or corporate name NOTE: Includes DMF original address or corporate name or change in DMF address or corporate name	1	1.3.1.1	Change of address or corporate name
	Change in contact/agent NOTE: Includes DMF original contact/agent or change in DMF contact/agent	1	1.3.1.2	Change in contact/agent
314.72	Change in ownership of an application	1	1.3.1.5	Change in ownership of an application
314.50(d)(1)(v)	Field copy certification	1	1.3.2	Field copy certification
Generic Drug Enforcement Act (GDEA)	Debarment certification	1	1.3.3	Debarment certification
314.94(13)	Financial certification and disclosure (Form FDA 3454 and Form FDA 3455)	1	1.3.4	Financial certification and disclosure

CFR Citation/Source		CTD /*STF Heading/**Attribute(s)			
NUMBER	TITLE	MODULE	NUMBER	TITLE	
314.50(h)	Patent information (Form FDA	1	1.3.5.1	Patent information	
314.53(e)	3542a and Form FDA 3542)				
314.94(12)	Patent certification	1	1.3.5.2	Patent certification	
314.95	Notice of certification of	1	1.3.5.3	Exclusivity claim	
	nonvalidity or noninfringement				
	of patent				
314.420(d)	Incorporating DMF information	1	1.4.1	Letter of authorization	
	by reference (authorization from				
	DMF holder)				
314.50(g)(1)	Written statement of	1	1.4.2	Statement of right of reference	
	authorization for references				
	(copy of LOA received from				
	DMF holders - submitted by				
214 420(1)	BLA, NDA, or IND applicants)	1	1.40		
314.420(d)	List of authorized persons to	1	1.4.3	List of authorized persons to incorporate	
21404(11)	incorporate by reference	1	1 4 4	by reference	
314.94(11)	Reference to information	1	1.4.4	Cross-reference to previously submitted	
214.65	previously submitted	1	1.5.5	information	
314.65	Withdrawal of an unapproved	1	1.5.5	Withdrawal of an unapproved BLA, NDA,	
214.150	application	1	1.5.6	ANDA or Supplement	
314.150	Withdrawal of listed drug	1	1.5.6	With drawal of listed drug	
314.150(c)	Request for withdrawal of	1	1.5.7	Withdrawal of approval of an application or revocation of license	
314.102	approval	1	1.6.1		
314.102	Communications: meetings	1	1.6.2	Meeting request Meeting background materials	
314.102	Communications: meetings	1	1.6.3	ž ž	
	Communications: meetings			Correspondence regarding meetings	
314.103(c)	Scientific and medical disputes	1	1.10.1	Request for dispute resolution	
314.103(c)	Scientific and medical disputes	1	1.10.2	Correspondence related to dispute resolution	
314.96	Amendment to an unapproved	1	1.11.1	Quality information amendment	
314.70	application: Chemistry	1	1.11.1	Quanty information amendment	
	application. Chemistry]		

CF	R Citation/Source	CTD /*STF Heading/**Attribute(s)			
NUMBER	TITLE	MODULE	NUMBER	TITLE	
	(information not fitting under Module 3)				
314.98	Amendment to an unapproved application: Toxicology (information not covered under Module 4)	1	1.11.2	Nonclinical information amendment	
314.96	Amendment to an unapproved application: Clinical (information not fitting under Module 5)	1	1.11.3	Clinical information amendment	
314.96	Multiple information amendment:	1	1.11.4	Multiple module information amendment	
	Request for comment and advice	1	1.12.4	Request for comments and advice	
GDEA	Generic drug enforcement act statement	1	1.12.10	Generic drug enforcement act statement	
314.94(a)(3)	Basis for abbreviated new drug application submission	1	1.12.11	ANDA basis for submission statement	
314.94(a)(4)	Conditions for use	1	1.12.11	ANDA basis for submission statement	
314.94(a)(5)	Active ingredient	1	1.12.12	Comparison of generic drug and reference listed drug	
314.94(a)(6)	Route of administration, dosage form, and strength	1	1.12.12	Comparison of generic drug and reference listed drug	
25.15(d)	Environmental impact analysis statement (if applicable)	1	1.12.14	Environmental analysis	
320.22 (a)	Request for waiver of in vivo bioavailability studies	1	1.12.15	Request for waiver of in-vivo bioavailability studies	
314.81(b)(i)(ii)	Field alert reports	1	1.12.16	Field alert reports	
314.81(b)(2)(i)	Annual Report: Summary	1	1.13.1	Summary of nonclinical studies	
314.81(b)(2)(i)	Annual Report: Summary	1	1.13.2	Summary of clinical pharmacology information	
314.81(b)(2)(i)	Annual Report: Summary	1	1.13.3	Summary of safety information	

CFR	Citation/Source	CTD /*STF Heading/**Attribute(s)			
NUMBER	TITLE	MODULE	NUMBER	TITLE	
314.81(b)(2)(i)	Annual Report: Summary	1	1.13.4	Summary of labeling changes	
314.81(b)(2)(i)	Annual Report: Summary	1	1.13.5	Summary of manufacturing changes	
314.81(b)(2)(i)	Annual Report: Summary	1	1.13.6	Summary of microbiological changes	
314.81(b)(2)(i)	Annual Report: Summary	1	1.13.7	Summary of other significant new information	
314.81(b)(2)(ii)	Annual Report: Distribution data	1	1.13.11	Distribution data	
314.81(b)(2)(vii)	Annual Report: Status report of clinical and nonclinical toxicology postmarketing study commitments	1	1.13.12	Status of postmarketing study commitments and requirements	
314.81(b)(2)(viii)	Status report of other (chemistry, manufacturing, controls) postmarketing study commitments	1	1.13.13	Status of other postmarketing studies and requirements	
314.81(b)(2)(ix)	Annual Report: Log of outstanding regulatory business	1	1.13.14	Log of outstanding regulatory business	
314.94(a)(8)(ii)	Copies of proposed labeling [Use appropriate sections]	1	1.14.1	Draft labeling	
314. 94(a)(8)(ii)	Draft carton and container labels	1	1.14.1.1	Draft carton and container labels	
314.50(c)(2)(i)	The proposed text of the labeling with annotations	1	1.14.1.2	Annotated draft labeling text	
314.94(a)(8)(ii)	Draft labeling text	1	1.14.1.3	Draft labeling text	
314.94(a)(8)(ii)	Final carton or container labels	1	1.14.2.1	Final carton or container labels	
314.94(a)(8)(ii)	Final package insert (package inserts, patient information, medication guides)	1	1.14.2.2	Final package insert (package inserts, patient information, medication guides)	
314.94(a)(8)(ii)	Final labeling text	1	1.14.2.3	Final labeling text	
314.94(a)(8)(iii)	Statement of proposed labeling	1	1.14.3.1	Annotated comparison with listed drug	
314.94(a)(8)(iv)	Comparison of approved and proposed labeling	1	1.14.3.1	Annotated comparison with listed drug	
314.94(a)(8)(i)	Listed drug labeling	1	1.14.3.2	Approved labeling text for listed drug	

CFR Citation/Source		CTD /*STF Heading/**Attribute(s)		
NUMBER	TITLE	MODULE	NUMBER	TITLE
314.94(a)(8)(i)	Labeling text for reference listed drug	1	1.14.3.3	Labeling text for reference listed drug
314.81(b)(3)(i)	Product labeling for 2253 submissions (if applicable)	1	1.14.6	Product labeling for 2253 submissions
202.1 314.81(b)(3)(i) 202.1(j)(4) 314.550 314.640	Regulations related to promotional materials [use appropriate sections]	1	1.15	Promotional material **[attribute = promotional-material-audience-type]
202.1 202.1(j)(4)	Request for advisory comments on launch materials	1	1.15.1.1	Request for advisory comments on launch materials
202.1 202.1(j)(4)	Request for advisory comments on non-launch materials	1	1.15.1.2	Request for advisory comments on non-launch materials
202.1 314.550	Presubmission of launch promotional materials for accelerated approval products	1	1.15.1.3	Presubmission of launch promotional materials for accelerated approval products
202.1 314.640	Presubmission of launch promotional materials for products approved when human efficacy studies are not ethical or feasible	1	1.15.1.3	Presubmission of launch promotional materials for accelerated approval products
202.1 314.550	Presubmission of non-launch promotional materials for accelerated approval products	1	1.15.1.4	Presubmission of non-launch promotional materials for accelerated approval products
314.640	Presubmission of non-launch promotional materials for products approved when human efficacy studies are not ethical or feasible	1	1.15.1.4	Presubmission of non-launch promotional materials for accelerated approval products

CFR Citation/Source		CTD /*STF Heading/**Attribute(s)		
NUMBER	TITLE	MODULE	NUMBER	TITLE
202.1 Section 503C of the Federal Food, Drug, and	Pre-dissemination review of television ads	1	1.15.1.5	Pre-dissemination review of television ads
Cosmetic Act 202.1	Response to untitled letter or warning letter	1	1.15.1.6	Response to untitled letter or warning letter
202.1	Response to information request	1	1.15.1.7	Response to information request
202.1 314.81(b)(3)(i) 202.1(j)(4) 314.550 314.640	Correspondence accompanying materials previously missing or rejected	1	1.15.1.8	Correspondence accompanying materials previously missing or rejected
202.1 314.81(b)(3)(i) 202.1(j)(4) 314.550 314.640	Withdrawal request	1	1.15.1.9	Withdrawal request
202.1 202.1(j)(4) 314.550 314.640	Submission of annotated references	1	1.15.1.10	Submission of annotated references
202.1	General correspondence	1	1.15.1.11	General correspondence
202.1 314.81(b)(3)(i) 202.1(j)(4) 314.550 314.640	Regulations related to submission of promotional materials [use appropriate sections]	1	1.15.2	Materials **[attribute = promotional-material-doc-type]
202.1 314.81(b)(3)(i) 202.1(j)(4) 314.550	Regulations related to promotional materials [use appropriate sections]	1	1.15.2.1	Material **[attributes =promotional-material-type, material-id, issue-date]

CFR Citation/Source		CTD /*STF Heading/**Attribute(s)		
NUMBER	TITLE	MODULE	NUMBER	TITLE
314.640				
202.1	Clean version	1	1.15.2.1.1	Clean version
314.81(b)(3)(i)				
202.1(j)(4)				
314.550				
314.640				
202.1	Annotated version	1	1.15.2.1.2	Annotated version
202.1(j)(4)				
314.550				
314.640				
202.1	Annotated labeling version	1	1.15.2.1.3	Annotated labeling version
202.1(j)(4)				
314.550				
314.640				
202.1	Annotated references	1	1.15.2.1.4	Annotated references
202.1(j)(4)				
314.550				
314.640				
FDAAA 505-1	Risk evaluation and mitigation	1	1.16	Use the appropriate sections
[355-1]	strategies (REMS)			
FDAAA	Correspondence regarding	1	1.17.1	Correspondence regarding postmarketing
	postmarketing commitments			commitments
FDAAA	Correspondence regarding	1	1.17.2	Correspondence regarding postmarketing
	postmarketing requirements			requirements
314.420(a)	Drug master files	1, 2, 3, 4, 5	As needed	Use appropriate sections
314.96	Amendment to an unapproved	3	As needed	Use appropriate sections
	application: Chemistry			
314.94(9)	Chemistry, manufacturing, and	3	As needed	Use appropriate sections
	control			
314.94(a)(7)	Bioequivalence	5	5.3	Use appropriate sections
314.96	Amendment to an unapproved	5	As needed	Use appropriate sections

ANDA Mapping Section

CFR Citation/Source		CTD /*STF Heading/**Attribute(s)		
NUMBER	TITLE	MODULE	NUMBER	TITLE
	application: Clinical			
314.94(a)(2)	Table of Contents	N/A	N/A	N/A

Appendix 2 – Module 1 Summary of Changes

A. Module 1 Summary of Changes (02/07/2014, version 2.3)

Module	Old Title	New Title
Section		
1.15.1.5	Promotional materials submitted pursuant to section 503B	Pre-dissemination review of television ads

B. Module 1 Summary of Changes (08/23/2013, version 2.2)

Module	Old Title	New Title
Section		
1.15.2.1	Material <attribute =="" [promotional-material-type]=""></attribute>	1.15.2.1 Material [attributes=promotional-material-type,
		material-id, issue-date]

C. Module 1 Summary of Changes (11/1/2012, version 2.1)

Module	Old Title	New Title
Section		
1.16	Risk evaluation and mitigation strategies (REMS)	Risk Management Plan
1.16.1	N/A	Risk Management (Non-REMS)
1.16.2	N/A	Risk Evaluation and Mitigation Strategy (REMS)
1.16.2.1	N/A	Final REMS
1.16.2.2	N/A	Draft REMS
1.16.2.3	N/A	REMS Assessment
1.16.2.4	N/A	REMS Assessment Methodology
1.16.2.5	N/A	REMS Correspondence
1.16.2.6	N/A	REMS Modification History

D. Module 1 Summary of Changes (6/1/2012, version 2.0)

Module	Old Title	New Title
Section		
1.1	Forms and form type e.g.	Forms
	1.1.1 Application form: FDA form 1571	Form ** [attribute = form-type]
	1.1.2 Application form: FDA form 356h	
	1.1.3 User fee cover sheet: FDA form 3397	
	1.1.4 Annual report transmittal: FDA form 2252	
	1.1.5 Advertisements and promotional labeling	
	transmittal: FDA form 2253	
	1.1.6 Transmittal of Labels and Circulars: FDA form 2567	
1.3.1.5	Change in ownership of an application	Change in ownership of an application or reissuance of
1.3.1.3	Change in ownership of an application	license
1.3.5.3	Exclusivity request	Exclusivity claim
1.3.6	N/A	Tropical disease priority review voucher
1.4.4	Cross reference to other applications	Cross-reference to previously submitted information
1.5.1	Withdrawal request	Withdrawal of an IND
1.5.5	Withdrawal of an unapproved NDA	Withdrawal of an unapproved BLA, NDA, ANDA, or
		supplement
1.5.7	Request for withdrawal of application approval	Withdrawal of approval of an application or revocation of
		license
1.7.4	N/A	Correspondence regarding fast track/rolling review
1.8.4.	N/A	Animal efficacy study for approval under the animal rule
1.9.5	Proposal for written agreement	No longer applicable
1.11.2	Safety information amendment	Nonclinical information amendment
1.11.3	Efficacy information amendment	Clinical information amendment
1.11.4	N/A	Multiple module information amendment
1.12.2	Request to charge	Request to charge for clinical trial
1.12.3	Notification of charging under treatment IND	Request to charge for expanded access

Module	Old Title	New Title
Section		
1.12.6	Exception from informed consent for research	Exception from informed consent for emergency research
1.12.7	Public disclosure statement for exception from informed consent for research	Public disclosure statement for exception from informed consent for emergency research
1.12.8	Correspondence regarding exception from informed consent for research	Correspondence regarding exception from informed consent for emergency research
1.12.11	Basis for submission statement	ANDA basis for submission statement
1.12.17	N/A	Orphan drug designation
1.13.12	Status of postmarketing study commitments	Status of postmarketing study commitments and requirements
1.13.13	Status of other postmarketing studies	Status of other postmarketing studies and requirements
1.13.15	N/A	Development safety update report (DSUR)
1.14.6	N/A	Product labeling for 2253 submissions
1.15	Promotional material	Promotional material <attribute =="" [promotional-material-audience-type]=""></attribute>
1.15.1	N/A	Correspondence relating to promotional materials
1.15.1.1	N/A	Request for advisory comments on launch materials
1.15.1.2	N/A	Request for advisory comments on non-launch materials
1.15.1.3	N/A	Presubmission of launch promotional materials for accelerated approval products
1.15.1.4	N/A	Presubmission of non-launch promotional materials for accelerated approval products
1.15.1.5	N/A	Promotional materials submitted pursuant to section 503B
1.15.1.6	N/A	Response to untitled letter or warning letter
1.15.1.7	N/A	Response to information request
1.15.1.8	N/A	Correspondence accompanying materials previously missing or rejected
1.15.1.9	N/A	Withdrawal request
1.15.1.10	N/A	Submission of annotated references
1.15.1.11	N/A	General correspondence
1.15.2	N/A	Materials <attribute =="" [promotional-material-doc-type]=""></attribute>

Module	Old Title	New Title
Section		
1.15.2.1	N/A	Material <attribute =="" [promotional-material-type]=""></attribute>
1.15.2.1.1	N/A	Clean version
1.15.2.1.2	N/A	Annotated version
1.15.2.1.3	N/A	Annotated labeling version
1.15.2.1.4	N/A	Annotated references
1.16	Risk management plans	Risk evaluation and mitigation strategies (REMS)
1.17	N/A	Postmarketing studies
1.17.1	N/A	Correspondence regarding postmarketing commitments
1.17.2	N/A	Correspondence regarding postmarketing requirements
1.18	N/A	Proprietary names
1.19	N/A	Pre-EUA and EUA
1.20	N/A	General investigational plan for initial IND